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(54) Title: BLENDS OF ELASTOMER BLOCK COPOLYMER AND ALIPHATIC ALPHA-OLEFIN/MONOVINYLIDENE ARO-MATIC MONOMER AND OR HINDERED ALIPHATIC VINYLIDENE MONOMER INTERPOLYMER

(57) Abstract

A thermopiastic elastomeric blend composition comprising: (A) from 99 to 1 weight percent of a styrenic block copolymer, and (B) from 1 to 99 weight percent of an interpolymer of (1) at least one aliphatic x-olenn and (2) at least one vinyildene aromatic monomer 97.33 a combination of at least one vinytidene aromatic monomer and at least one nindered aliphatic vinytidene monomer, wherein said vinyindene promatic monomer is present in an amount of from 0.5 to (5 mole % or from 17 to 55 mole % in said interpolymer, and

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BLENDS OF ELACTOMER BLOCK COPOLYMER AND ALIPHATIC ALPHA-CLEFIN MONOVINYLIDENE AROMATIC MONOMER AND/OF HINDERED ALIPHATIC VINYLIDENE MONOMER INTERPOLYMER

This invention relates to a thermoplastic elastomeric blend composition comprising an elastomer clock copolymer and an $\alpha-$ plefin/monovinylidene aromatic monomer and/or nindered aliphatic vinylidene monomer interpolymer, and to fabricated articles made therefrom.

used in the industry, frequently for impact modification of thermoplastic resins and engineering thermoplastics or for compatibilization of different types of resins. The class of elastomer saturated rupper plock copolymers, such as styrene-ethylene/putene-styrene plock copolymers 3-EB-S, however, is expensive to produce and difficult to process. The class of unsaturated rupper plock copolymers such as Styrene-Butadiene-Styrene (SBS) is susceptable to degradation. Blends of polymer components not readily miscible with these plock copolymers, such as conventional polyethylenes, can exhibit inferior mechanical properties, especially elastomeric retention such as permanent tensile set after elongation.

It would be desirable to provide either lower cost or more durable blends of these block copolymers by adding a polymeric component that does not significantly decrease the required performance properties. It would be further desirable to provide blends of these block copolymers and a polymeric component thereby maintaining or improving the performance of these block copolymers while providing low maze blends. It would also be desirable to maintain a low value for hardness as desired for applications requiring flexibility such as sheet, film or tubing. It would also be 30 desirable to maintain a high level of ultimate tensile properties to enhance the strength properties of the blends. Finally it would also be desirable to maintain a high level of % stress relaxation to enhance the surface conformity of the blends especially in 35 applications requiring covering of a surface.

According to the present invention there is provided a thermoplastic elastomeric blend composition comprising:

- (A) from 99 to 1 weight percent of a styrenic block copolymer; and
- 40 (B) from 1 to 99 weight percent of an interpolymer of (1) at least one aliphatic α -olefin and (2) at least one vinylidene aromatic

monomer or op a simplication of at least one vinylidene aromatic monomer and at least one numbered allicatio convildene monomer, wherein said vinvildene aromatic monomer to present in an amount if from ... To 10 mole or from ... To 48 mole on said interpolymer.

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In yet another under, the present invention browness $\hat{\gamma}$ tabricated article made from subn a thermoplastic quastometric blend composition.

In yet another appect, the present invention comprises an adhesive or adhesive commutation or a sealant formulation containing the aforementioned plends.

The term "interpolymer" is used herein to indicate a polymer wherein at least two different monomers are polymerized to make the interpolymer.

The term "hydrocarpyi" means any aliphatic, symboliphatic, aromatic, aryl substituted aliphatic, aryl substituted symboliphatic, aliphatic substituted aromatic, in symboliphatic substituted aromatic groups. The aliphatic or cyclosliphatic groups are preferably saturated. Likewise, the term "hydrocarpyloxy" means a hydrocarpyl group having an oxygen linkage between it and the carbon atom to which it is attached.

The term "monomer residue" means that portion of the polymerizable monomer molecule which resides in the polymer chain as a result of being polymerized with another polymerizable molecule to make the polymer chain.

The term "substantially random" in the substantially random interpolymer comprising an α -olefin and a vinylidene aromatic monomer or hindered aliphatic vinylidene monomer as used herein means that the distribution of the monomers of said interpolymer can be described by the Bernoulli statistical model or by a first or second order Markovian statistical model, as described by J. C. Randall in <u>POLYMER SEQUENCE DETERMINATION</u>, Carbon-13 NMR Method, Academic Press New York, 1977, pp. 71-78. Preferably, the substantially random interpolymer comprising an α -olefin and a vinylidene aromatic monomer does not contain more than 15 percent of the total amount of vinylidene aromatic monomer in clocks of vinylidene aromatic monomer in more than 3 units. More preferably, the interpolymer was not characterized by a high degree of either isotacticity or syndiotacticity. This means that in the carbon-13 NMR spectrum of the substantially random interpolymer the peak areas corresponding to the main chain methylene

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and methane (arbono numberentin'i estado des) mad dequencts in pasemio mad equences unotad numberodo "" percent un the foral peak area or the main (num mothylene and pethane) farbons.

The term office untigment of theoretic to mean elastomers caving at least the office removed that a dark proper consider into dark property and at least one place removed that the term of the place of the linear of radial type, wither diploce of the term of the terms.

Any numerical willed resisted herein include it wildes from the lower value to the upper walde in indicements it into init provided that there is a deparation of a least of units between any lower value and any higher value. As an example, if it is stated that the amount of a somponent or a value of a process variable such as, for example, temperature, pressure, time and the like is, for example, from 1 to 30, preferably from 20 to 60, more preferably from 30 to 70, it is intended that values such as 15 to 65, 20 to 68, 43 to 81, 30 to 32 etc. are expressly enumerated in this specification. For values which are less than one, one unit is considered to be 0.0001, 1.001, 0.01 or 0.1 as appropriate. These are only examples of what is specifically intended and all possible symboliations of numerical values between the lowest value and the mignest value enumerated are to be considered to be expressly stated in this application in a similar manner.

Suitable unsaturated block copolymers include those represented by the following formulas:

Formula I $A-B-R(-B-A)_n$ or

Formula II $A_x = (BA +)_y - BA$

wherein each A is a polymer block comprising a monovinylidene aromatic monomer, preferably styrene, and each B is a polymer block comprising a conjugated diene, preferably isoprene or butadiene, and optionally a monovinylidene aromatic monomer, preferably styrene; E is the remnant of a multifunctional soupling agent; n is an integer from 1 to 5; x is zero or 1; and y is a real number from zero to 4.

The preparation of the plock dopolymers useful herein is not the subject of the present invention. Methods for the preparation of such

whose sapolymens (19) as we in the art. Sostable outalysts for the preparation i isetui ullok oppolomero with unsaturated supper monomer inits include lithium rased ratalysts and especially lithium-alkyrs. T.C. Tat. No. 1,575, His describes suitable methods for hydrogenation or block a polymena with andaparate; rupper a domei unito to from Clock Currymers with Churated supper schames inits. The structure if the polymers is retermined by their methods of polymerication. For example, linear polymers result by sequential introduction of the resired rupper monomer into the relation vesse, when coing such unitiators as lithium-alayes or milithiostilbene and the like, or by soupling a two seament block sopolymer with a difunctional soupling agent. Branched structures, in the other hand, may be obtained by the use of suitable soupling agents having a functionality with respect to the block copolymers ...th unsaturated rupper monomer inits of three or more. Coupling may re-effected with multifunctional coupling agents such as dihalogikanes in alkenes and divinyl benzene as well as with sertain polar compounds such as silicon halides, silomanes or esters of mononyaria alcoholo with sarboxylia saids. The presence of any coupling residues in the polymer may be ignored for an adequate description of the block topolymers forming a part of the composition of this invention.

Suitable block sopolymers having unsaturated rubber monomer units includes, but is not limited to, styrene-butadiene (SB), styrene-isoprene(SI), styrene-butadiene-styrene (SBS), styrene-isoprene-styrene (SIS), α -methylstyrene-butadiene- α -methylstyrene and α -methylstyrene-isoprene- α -methylstyrene.

The styrenic portion of the block copolymer is preferably a polymer or interpolymer of styrene and its analogs and nomologs including α -methylstyrene and ring-substituted styrenes, particularly ring-methylated styrenes. The preferred styrenics are styrene and α -methylstyrene, and styrene is particularly preferred.

Block copolymers with unsaturated rubber monomer units may comprise homopolymers of butadiene or isoprene or they may comprise copolymers of one or both of these two dienes with a minor amount of styrenic monomer.

Preferred block sopolymers with saturated rupper monomer units comprise at least one segment of a styrenic unit and at least one segment of an ethylene-putene or ethylene-propylene copolymer. Preferred examples of outpublick copolymers with saturated rupper monomer units include styrene/ethylene-putene copolymers.

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Tyrens ethylohemoropylohe oblolomero, hurene ethylenemotyrene otyrene obspolomero, hyrene ethylenemopylene otyrene obspolomero.

Appropriation of plack supplyments with unsaturated rupper someoner units is projectedly different by see if a saturated supprising the reaction products of an eluminum alays sembound with nickel of conalt surpoxylates is alkowinged under such conditions as to substantially sampletely hydrogenate at least to percent of the alignatic souple bonds while hydrogenating no more than is percent of the styrenis aromatic souple bonds. Preferred plock supplyments are noted where at least to percent if the alignatic double bonds are hydrogenated while less than a percent of the aromatic souple bonds are hydrogenated.

The proportion of the objection and denerally between and so as percent by weight in the total swight of the plack objective. Ereferably, the block objectivens contain from 1, to 35 weight bendent of objectiveness block segments and from 40 to 45 weight percent of rubber monomer block segments, based in the rotal weight of the block copolymer.

The average molecular weights of the individual blocks may vary within certain limits. In most instances, the styrenic block segments will have number average molecular weights in the range of 5,000 to 125,000, preferably from 7,000 to 60,000 while the rubber monomer block segments will have average molecular weights in the range of 10,000 to 300,000, preferably from 30,000 to 100,000. The total average molecular weight if the block sepolymer is typically in the range of 25,000 to 250,000, preferably from 35,000 to 200,000.

Further, the various block copolymers suitable for use in the present invention may be modified by graft incorporation of minor amounts of functional groups, such as, for example, maleic anhydride by any of the methods well known in the art.

Block appolymers useful in the present invention are sommercially available, such as, for example, supplied by Chell Chemical Company under the designation of MRATONIM and supplied by Dexco Polymers inder the designation of VECTORIM.

Blend component (B) for the polymer stend composition of the present invention include, but are not limited to, substantially random interpolymers prepared by polymerizing one or more α -olefin monomers with one or more vinylidene aromatic monomers and/or one or

more hinder-H alibhatic in SypicalSphatic vinylidene monomers, and phionally with other colymericable ethylenibully unbaturated monomers.

deltable α - ref. numbered contained in the blend component β include it example alignatic and systematic α -obtains having from β to is carpon example, and preferably α -prefixe having from β to β carpon atoms. Most preferably, the alignatic α -obefix of blend component β , comprises ethylene or propylene, preferably ethylene, optionally together with one or more other α -obefine having from β carpon atoms, outhly as ethylene and propylene, or ethylene and obtene, or ethylene and propylene, or ethylene and

Suitable vinylidene iromatic monomers which can be employed to prepare the interpolymers include, for example, those represented by the following cormula:

$$\begin{array}{c}
Ar \\
(CH_2)_n \\
 & (CH_2)_n
\end{array}$$

$$R^1 - C = C(R^2)_2$$

wherein \mathbb{R}^1 is selected from the group of radicals consisting of hydrogen and alkyl radicals containing from 1 to 4 carbon atoms, preferably hydrogen or methyl; each \mathbb{R}^1 is independently selected from the group of radicals consisting of hydrogen and alkyl radicals containing from 1 to 4 carbon atoms, preferably hydrogen or methyl; Ar is a pnenyl group or a pnenyl group substituted with from 1 to 5 substituents selected from the group consisting of halo, $\mathbb{C}_{\mathbb{R}^d}$ -alkyl, and $\mathbb{C}_{\mathbb{R}^d}$ -haloalkyl; and n has a value from zero to 4, preferably from zero to 2, most preferably zero. Exemplary monovinylidene aromatic monomers include styrene, vinyl toluene, α -methylstyrene, t-butyl styrene, chlorostyrene, including all isomers of these compounds. Particularly suitable such monomers include styrene and lower alkylor halogen-substituted derivatives thereof. Preferred monomers include styrene, a-methyl styrene, the lower alkyl- $(\mathbb{C}_1 - \mathbb{C}_4)$ or pnenyl-ring substituted derivatives of styrene, such as for example,

By the term "hindered aliphatic or sycloaliphatic vinylidene compounds", it is meant addition polymericable vinylidene monomers corresponding to the formula:

monovinylidene monomer is styrene.

ortho-, meta-, and para-methylstyrene, the ring nalogenated styrenes, para-vinyl toluene or mixtures thereof. A more preferred aromatic

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$$R^1 - C = C(R^2)$$

wherein Artiff a literically colky. Galphatic or cycloalignatic supstituent of apt. Suppond, we is something from the group of radicals observed or symposes and alays produced bintaining from 1 5 to 4 harpon store, is resumbly hydroden is detayl; wach Prize independently released from the group of radicald constating of hydrogen and alkyl susteals sontaining from 1 to 4 sarbon atoms, preferably hydroden is methyl, in alternatively Et and At together form a ring system. By the term "steriogally bulky" is meant that the monomer bearing this capatituent is normally incapable or addition polymerization by standard Cledier-Matta polymerization sacalysts at a rate comparable with the terred bolymerications. Scenerical nindered aliphatic of sycloarinhatic minuliaens compounds lie monomers in which the of the dirpon stand agaring othylenic andapuration of terplary or alipnatic groups duon as hyphonexy., syclonexenyl, syclocotenyl, or ring alkyl or aryl substituted derivatives thereof, tert-butyl, norbornyl. Most preferred hindered aliphatic or cycloaliphatic vinylidene compounds are the warious icomeric vinyl- ring substituted derivatives of dyclonexene and substituted dyclonexenes, and 5ethylidene-3-norpornene. Especially suitable are 1-, 3-, and 4vinyloyolshexene.

The interpolymens of one in more α -claims and one or more monovinylidene aromatic monomers and/or one or more nindered alighatic or syctoaliphatic winviluene monomers employed in the present invention as component. By are substantially random polymers. These interpolymers usually contain from 1 to 65, preferably from 5 to 60, more preferably from 10 to 55 mole percent of at least one winylidene aromatic monomer and/or nindered alighatic or cycloaliphatic winylidene monomer and from 35 to 99, preferably from 40 to 95, more preferably from 45 to 30 mole percent of at least one alighatic α -olefin having from 2 to 20 carpon atoms

Preferably, higher molecular weight interpolymer component (B) are used, such as those that possess a number average Mw of greater than 13,000. Also preferably such polymers possess a melt index (I), ASTM D-1238 Procedure A. condition E, or less than 125, more preferably from 1.31 - 100, even more preferably from 0.31 to 25, and most preferably from 0.15 to 0. Also, the substantially random interpolymers have a heat of fusion of less than 50 Jog.

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The present invention provides clends of interpolymer components of molecular weight and composition instributions selected to obtain an overall molecular weight and composition instribution which lives enhanced properties or processability.

While preparing the oupstantially random interpolymers, component [8], is will be described hereinarths, in amount of stactic vinylidene aromatic nomopolymer may be formed and to nomopolymerization in the vinylidene arcmatic monomer at elevated temperatures. In general, the nigher the polymerication temperature was, the nigher is the smount of homopolymer cormed. The presence of vinylidene aromatic nomopolymer is in general not detrimental for the purposes of the present invention and may be tolerated. The vinylidene aromatic nemopolymer may be separated from the interpolymer, if desired, by extraction techniques such is selective predipitation from Jolution with a non scient for either the interpolymer in the vinylidene iromatic nomopolymer. For the purpose of the present invention it is preferred that no more than 20 weight percent, preferably less than is weight percent based on the total weight of the interpolymers of vinylidene aromatic nomopolymer is 20 present.

The substantially random interpolymers may be modified by typical grafting, hydrogenation, functionalizing, or other reactions well known to those skilled in the art. The polymers may be readily sulfonated or chlorinated to provide functionalized derivatives according to established techniques.

The substantially random interpolymers are prepared by polymerizing a mixture of polymerizable monomers in the presence of metallocene or constrained geometry catalysts.

described in US Application Serial Number 545,403 filed July 3, 1990 (corresponding to EP-A-0,416,815) by James C. Stevens et al., both of which are incorporated herein by reference in their entirety. Preferred operating conditions for such polymerization reactions are pressures from atmospheric up to 3000 atmospheres and temperatures from -30°C to 200°C. Polymerizations and unreacted monomer removal at temperatures above the autopolymerization temperature of the respective monomers may result in formation is some amounts of homopolymer polymerization products resulting from free radical polymerization.

The substantially random of leafin vanyindene aromatic interpolymers can also be prepared by the methods seconded by John 9. Bradfute et al. W. F. Grade v 7: W 46 80096; by R. B. Pannell Exxon Shemicul Latents, Inc. ... w 41 0807; and in <u>Clastics Technology</u>, b. 07 September 1990, ... b which are incorporated nerein by reference in their entirety.

The substantially random α_{τ} letter winviliagene aromatic interpolymens can also be prepared by the methods described in JP 37/278230 employing compounds shown by the general formula

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where Cp^* and Cp^* are hydrogentialienyl groups, indenyl groups, cluorenyl groups, in substituents of these, independently if each other; P^* and P^* are hydrogen tims, whilsden atoms, hydrogarbon groups with carbon numbers of 1-12, alkowyl groups, in anylowyl groups, independently of each other; M is a group IV metal, preferably Cp^* or Hf, most preferably Cp^* and R^* is an alkylene group or silanediyl group used to cross-link Cp^* and Cp^*).

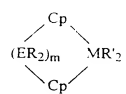
Also suitable are the substantially random interpolymers which possess at least one α -olefin/vinyl aromatic/vinyl aromatic/ α -olefin tetrad disclosed in a copending application by Francis J. Timmers et al. filed on the same date as this application. These interpolymers contain additional signals with intensities breater than three times the peak to peak noise. These signals appear in the chemical shift range 43.70-44.25 ppm and 38.0-38.8 ppm. Specifically, major peaks are observed at 44.1, 43.9 and 38.2 ppm. A proton test MMR experiment indicates that the signals in the chemical shift region 43.70-44.25

ppm are methyles (arbons and the signals in the region (4.)-(5.) ppm are methylene (arbons.

in order to determine the farbine. DMR membed united of these interpolyment, the full wind princedured in conditions are employed. A five to ten estant jerbent polyment of interpolation to prepared in a mixture consisting to the construction of the construction of the modern arrowmium trip (abetylapetonate) in 1,2,4-triphloropenses. TMR spectra are acquired at 130°D using an inverse gated decoupling beguence, a 90° pulse width and a pulse selay of five seconds or more. The spectra are referenced to the instanted methylene signal of the polymer assigned at 30,000 ppm.

It is believed that these new signals are due to dequences involving two dead-fit-tail winyl arcmatic monomer preceded and followed by at least the α -disfiningertion, for example, an ethylene/styrene/styrene/styrene tetradwherein the styrene monomer insertions in said tetrads occur exclusively in a l,w head to tail) manner. It is understood by one skilled in the art that for such tetrads involving a winyl aromatic monomer other than styrene and an α -diefin other than ethylene that the ethylene/winyl aromatic monomer/vinyl aromatic monomer/ethylene tetrad will give rise to similar carbon-13 MMR peaks but with slightly different chemical shifts.

These interpolymers are prepared by conducting the polymerization at temperatures of from -30°C to 250°C in the presence of such catalysts as those represented by the formula



wherein: each Cp is independently, each occurrence, a substituted cyclopentadienyl group π-bound to M; E is C or Si; M is a group IV metal, preferably 2r or Hf, most preferably 2r; each R is independently, each occurrence, H, hydrocarbyl, silahydrocarbyl, or hydrocarbylsilyl, sontaining up to 30 preferably from 1 to 20 more preferably from 1 to 10 darbon or silicon atoms; each R' is independently, each occurrence, H, halo, hydrocarbyl, hydrocarbyl, hydrocarbylsilyl containing up to 31 preferably from 1 to 20 more preferably from 1 to 10 darbon or silicon atoms or two P' groups together can be a C₁₋₁₀ hydrocarbyl substituted 1,3-butadiene; m

id longer and entirely, but sintending in the presence of an approvating repartatyon. Cartifularly, caltable dispetituted representational fround include the clustrates by the formula:

anderein each P lu independently, wear indurrence, H, hydrodarbyl, dilunyarodarbyl, in hydrodarbyldlyl, duntalning up to 30 preferably from 1 to 20 more preferably from 1 to 17 darbon in dilicon atoms or two R groups together form a divalent derivative if such group. Preferably, P independently each indurrence is unadiaging where appropriate all loomers invarided, methyl, stoyl, dropyl, butyl, pentyl, nexyl, denzyl, dhenyl is dilyl in where appropriate) two such a groups are linked i dether filming a fused find dystem such as indenyl, ilustenyl, Schringgroupskingenyl, retrangardilustenyl, in openyl, detrangardilustenyl, in openyl, detrangardilustenyl, in

Particularly preferred satalysts include, for example, rademic-dimethylsilanedlyl.d-methyl-4-phenylindenyl-Carconium dichloride, racemic-(dimethylsilanedlyl C-methyl-4-phenylindenyl-Carconium 1,4-diphenyl-1,3-butadiene, racemic-dimethylsilanedlyl(C-methyl-4-phenylindenyl) sirconium di-C_{1,4} alkyl, racemic-dimethylsilanedlyl(C-methyl-4-phenylindenyl) sirconium di-C_{1,4} alkyl, racemic-dimethylsilanedlyl(C-methyl-4-phenylindenyl) sirconium di-C_{1,4} alkoxide, or any combination thereof. Also included are the titanium-based satalysts, (N-(1,1-simethylethyl)-1,1-dimethyl-1-1,1,3,4,5-η)-1,5,6,7-tetranydro-s-indacen-1-yl;silanaminats a--B,titanium dimethyl; l-indenyl) (tertoutylamido) dimethyl- ilane ritanium dimethyl; l-indenyl) (tertoutylamido) dimethyl; and (3-isc-propyl),1,2,3,4,5-η)-1-indenyl) (tertoutyl amido) dimethylsilane titanium dimethyl, or any combination thereof.

B) of the present invention have been described in the literature.
Longo and Grassi (Makromol. Them., Volume 191, pages 1387 to 2396 (1990)) and D'Anniello et al. Journal of Applied Polymer Edience,
Volume 59, pages 1701-1704 (1995) reported the use of a catalytic system based on methylalumoxane (MAO) and systemeasurenyltitanium trichloride (CpTiCl. to prepare an ethylene-styrene sopolymer. Mu and Lin (Polymer Preprints, Am.Chem.Scs.,Civ. Polym. Them., Volume 35, pages 686,687 (1994) have reported sopolymerization using a TiCl.,NdCl.,Al(iBu), Matalyst to give random sopolymers of styrene and

pages 1453 (1.140 1931) have rescribed the appolymerization of conviewe and styrene within a T. 71 MdCl MdCl Al.Et a lataryst. The manufacture of devices in these objects monomer intercolymers auon as cropylene styrene and strene objects the described in Shited States hatent number (1.14.3%), industries that All the above methods industries of all the above methods industries along apponents are incorporated herein by reference.

Generally, the blend composition of the present invention comprises from 1 percent () 33 percent, by weight of the composition, 10 of the substantially random interpolymer of alighatic α_{τ} clefin/vinylidene aromatic monomer component (\mathbb{R}) and from 99 to 1percent, by weight of the composition, of the elastomer block copolymer component A.. Freierably, the composition comprises from 1 percent to 50 percent, more preferably from 10 to 45 percent, by weight of the composition, if the interpolymer of aliphatic α plefin/vinylidene arcmatic monomer component B) and from 00 to 50 percent, more preferably from 90 to 55 percent, by weight of the composition, of the elastomer block copolymer component (Λ) . For the sempositions containing at the most 50 weight percent of interpolymer 20 of aliphatic $\alpha\text{-olefin/vinylidene}$ aromatic monomer component (B), the compositions maintain a reasonably low Shore A hardness (65 or lower) as well as a fairly constant peak tonsile stress, which are desirable properties for elastomeric materials.

For the blend compositions in which the monovinylidene aromatic monomer component of the disphatic α -defin/monovinylidene aromatic interpolymer component to present in an amount of amount of from 17 to 65 mole 3, preferably from 20 to 65 mole 3 in said interpolymer, and fabricated articles made from such composition and said interpolymer is present in the blend at a concentration of 25 wt%, then the % stress relaxation of the blends is $\geq 38\%$. Further for the blend compositions in which the monovinylidene aromatic monomer component of the aliphatic α -olefin/monovinylidene aromatic interpolymer component is present in an amount of from 17 to 65 mole 1, preferably in an amount of from 20 to 65 mole 1 and said interpolymer is present in the blend at a concentration of 75 wt%, then the 1 stress relaxation of the blends is $\geq 65\%$.

Also for the blend compositions in which the monovinylidene aromatic monomer component of the alignatic α -elefin, monovinylidene aromatic interpolymer component is present in an amount of from 0.5 to 15 mol3 and said interpolymer is present in the blend at a

Vancentruition of Court Court Court Court of the plends is 2 11 MFs. Cortans is a constitution of the plends is 2 monthly court of the stimute of the stimut

Additives duch as inticxidants for example, hindered phenolics for example, Irdanox- 1010, phosphites for example, Irdanox- 1060), plind additives for example, FIB, inticlical additives, colourants, pigments, fillers, and the like our also se incrused in the present compositions, to the extent that they is not interiere with the enhances properties indicavers as a applicants.

The compositions of the present invention are compounded by any convenient mithous, including my clending the individual components and subsequently melb mixing, either directly in the extruder or mill used to make the conjuned article for example, the automotive part), or by pre-melb mixing in a separate extruder or mill for example, a Banbury mixer.

There are many types in molding iperations which can be used to form useful fabricated articles or parts from the present compositions, including various inhection molding processes (for example, that described in Modern Plastics Encyclopedia/89, Mid October 1988 Issue, Volume PI, Number 11, pp. 264-268, "Introduction to Injection Molding" and in pp. 277-271, "Injection Molding Thermoplastics", the discribures of which are incorporated herein by references and blow molding processes for example, that described in Modern Plastics Encyclopedia/89, Mid October 1988 Issue, Volume 65, Number 11, pp. 217-218, "Extrusion-Blow Molding", the disclosure of which is incorporated herein by reference) and profile extrusion. Some of the fabricated articles include sports articles, containers such as for food or other nousencid articles, footware, automotive articles, such as soft facia, sealants and assembly adhesives.

The compositions of the present invention can be further combined with many thermoplastic polymers to improve their properties, such as impact properties.

The compositions it the present invention are useful as tubing, tapes, adhesives, film, maskets. The compositions of the present invention are especially useful for the preparation of erastic films and fibres, asphalt blends, wire and cable, cound parriers, wax

plends, response senditive singular and in the impart modification of thermoplestics.

The inventi bouill be firth rollinitated for means of the collewing examples without limits for the inventi bothereps.

The respect of a state of the continuous continuous recommends.

Tensity to provided by the manufacturer of optained by ADTM 0 192.

With Percent Covrene as optained by 35 MMR.

The icllowing components are employed in the Examples and Comparative Experiments.

GENERAL PROCEDURE A FOR PREPARATION OF ETHYLENE/STYRENE COPOLYMERS A-E
Polymer to prepared in a 1 dallon upitated demi-continuous patch
reactor. The reaction mixture denerally consisted of a solvent
comprising systemexane or loopar-ET, and styrene. Frior to addition,

- colvent, Utyrene and Othylene are purified to remove water and oxygen.

 The inhibitor in the Chyrene is also removed. Inerts are removed by purging the vessel with ethylene. The vessel is then pressure
- controlled to a set point with ethylene. Hydrogen is added by filling a 70 cd cylinder to a set pressure and then adding it to the reactor to control molecular weight. Temperature in the vessel is controlled
- to set-point by varying the cooling flow of the cooling coals within the vessel. Prior to polymerization, the vessel is heated to the desired run temperature and the catalyst components: Titanium (N-1,1-dimethylethy) dimethyl 1-(1,2,3,4,5-n)-2,3,4,5,-tetramethyl- 2,4-syclopentadien-)-yl.gilanaminato: [1-(N)-dimethyl, [JAS#135072-62-7,
- Tris(pentafluorophenyl/poron, JAS#J71103-15-5, Modified methylaluminoxane Type JA, JAS# 146905-79-5, are combined in mole ratios as shown and added to the vessel. After starting, the polymerization is allowed to proceed with ethylene supplied to the reactor as required to maintain vessel pressure. In some cases,
- hydrogen is added to the headspace of the reactor to maintain a mole ratio with respect to the ethylene concentration. At the end of the run, about 1,000 ppm or TrganoxTM 1010 anti-oxidant is then added to the solution on a polymer basis and the polymer is isolated from the solution by precipitation with methanol. The resulting polymers are
- 35 dried in a vacuum oven. The following Table 1 contains the reactor conditions.

Table 1

Namber Oambie		_ Weitt	twene		(r. 1991. 100 (estur estur	nu- 'acre' 'ame'	-,,mer
] amu	<u> </u>	ime	131	·. E .	1 1152	, n.s.3		rams
E .7 - ₃ ,		.Jonar : BE	,		1,10		47.234		i
E 3	1122	l dydiue nexame	:11	1 : . 3	,				
E. 3 -	1-1.	-Supar #E	-10	1	-, - =				115
E/3 - 1	* # Ī	luopar -	-,	372	-7.52		- : :	1:0	:52
E. J 4		1 3 A		 	1 7 7 7	1 . !		100	

Table 1 contd.

Jumbig Number	Prin Prink Prink	Tyrene Tyrene Tyrene Tyrer	TI B A1	Efficiency Typiyar
<u> 변 . </u>	<u> </u>			
5// - E		:		1 - 1
E. ∪ =		44.1	1/3/11	1 200,000
E/C - 10 1	E. •			
E//		=	-	1200,200

* 190°C/2.2 kg

GENERAL PROCEDURE B FOR PREPARATION OF ETHYLENE/STYRENE COPOLYMERS F-H

5 EC1 Catalyst dimethyl(N-vi,l-dimethylethyl -1,l-dimethyl-i-[11,2,3,4,5-nv-1,5,4,1-retranydro-v-phenyl-v-indagen-iyl]silanaminato(Z-v-N-vitanium) Freparation.

Preparation of 3.4, ϵ , 7-February drop-shipdring a centle 2H shope. Indan 44.00 4, .7354 moles and 3-unleropropiony: enloride :0 100.99 q, 0.7354 moles) were stirred in CHCl. 300 mL, at 0°C as AlCl, (130.00 g, 0.3750 moles) was added slowly under a nitrogen flow. The mixture was then allowed to stir at room temperature for 2 hours. The volatiles were then removed. The mixture was then cooled to 0°C and concentrated $\mathrm{H}_2\mathrm{SO}_4$ (500 mL) slowly added. The forming solid had to be frequently broken up with a spatula as stirring was lost early in this step. The mixture was then left under nitrogen overnight at room temperature. The mixture was then heated until the temperature readings reached 90°C. These conditions were maintained for a 2 hour period of time during which a spatula was periodically used to stir the 20 mixture. After the reaction period crushed ice was placed in the mixture and moved around. The mixture was then transferred to a peaker and washed intermittently with $\mathrm{H}_2\mathsf{C}$ and diethylether and then the fractions filtered and combined. The mixture was washed with $\rm H_2O$ (2 $\rm x$ 200 mL). The organic layer was then separated and the volatiles removed. The desired product was then isolated via recrystallization from hexane at $C^{\circ}C$ as pale yellow crystals (22.36 g, 16.3% yield). H NMR (CDC1,): d2.04-2.19 (m, 1 H), 1.65 f, PHH=5.7 Hd, 1 H), 2.84-3.0 $_{\odot}$ m, 4 H , 3.03 $_{\odot}$ t, $^{\circ}$ [$_{\mathrm{HH}}$ =5.5 Hz, $_{\odot}$ H), 7.26 G, 1 H), 7.23 s, 1 30 H). ''S NMR | SDG1. : 425.71, L6.31, 32.1-, 33.84, 6.93, 118.40, 122.10, 135.88, 144.06, 152.84, 154.86, 106.50. 30-MS: Daloulated for _{1,4}H₁₂O 172.39, found 172.05.

35 Freparation of 1,2,3,.-Tetranydro-Tepnenyl-s-indaden.

Notes was ottered in description of and 10.10 in 100967 Notes was ottered in description of action was added clower. This mixture was onen allowed to the evention at room remperature. After the respect of period the mixture was duentied by fourth over the five mixture was onen actioned action period the mixture was duentied by fourth over the five mixture was onen actional five period with more five mixture was onen actional five with HOL and other was onen departed and washed with Holl and then order interesting and washed by the removal of the volatiles resulted in the isolation of the desired product as a tark oil 14.66 g, vi.3. Wield:

HOME FIGURE 12.0-1.1 m, 2.87, 1.6-3.1 m, 4.8 , vi.54 m, 18), 7.0-7.6 m, 7.83.

SCHMS: Valculated for J₁₆H₁, 232.13, cound 232.35.

Treparation of 1,2,2, -Twtranyard-T-phenvi- -indadene, illithium sait.

1,2,1,1=Tetranvari-T-phenvi-s-indaden 11.05 %, 1020 moles
was othered in nexame 150 min as nBuLt 1.36% moles, 10.00 mL of 2.0
M solution in systemewans was slowly added. This mixture was then allowed to stir overnight. After the reaction period the solid was sollected via suction filtration as a yellow solid which was washed with hexame, dried under vacuum, and used without further purification or analysis 12.2075 g, 31.10 yield).

Preparation of Chlorodimethyl(1,8,6,7-tetrahydro-3-phenyl-s-indaden-1-yl)silane.

25 1,2,3,5-Tetranydro-7-pnenyl-s-indacene, dilithium salt (12.2075 g, 0.05102 moles) in THF (50 mL) was added dropwise to a solution of Me-SiCl (3.5010 g, 0.1511 moles: in THF 100 mL) at 0.0. This mixture was then allowed to other at room temperature overnight. After the reaction period the volatiles were removed and the residue extracted and filtered using nexame. The removal if the hexame resulted in the isolation of the desired product as a yellow oil (15.1492 g, 91.10 yield). 1 H NMR (CDC1 $_{2}$): $_{2}$ 30.33 s, 3 H), 0.38 (s, 3 H), 2.20 'p, $_{3}$ 3 $_{
m HH}$ =7.5 Hz, 2 H), 2.9-3.1 (m, 4 H), 3.84 (s, 1 H), 6.69 (d, 3)HH=2.8 Hz, 1 H), 7.3-7.6 (m, 7 H), 7.68 (d, $^3J_{\rm HH}$ =7.4 Hz, 2 H). ¹³C NMR (CDC1, : d0.24, 0.38, 06.28, 33.05, 33.18, 46.13, 116.42, 119.71, 127.81, 128.88, 128.84, 129.86, 136.51, 141.81, 141.86, 142.17, 142.41, 144.62. GC-MS: Calculated for JLHLC101 324.11, found 324.38. 40

Freparation of N- \cdot 1,1-Dimetrylethyl.-1,1-dimetryi-1-1,5,6,7-tetrahydro-3-phenyl-s-indacen-1-yl)silanamine.

H NMR LIDG1, 1 40.02 5, 3 4), 0.04 5, 3 H), 1.27 5, 9 H), 2.16 (p, 0.05 Hz, 2 Hz, 2 H), 2.4-3.0 m, 4 H), 3.68 (s, 1 H), 6.69 (s, 1 H), 7.3-7.5 (m, 4 H), 0.63 (d, 0.04 Hz, 2 H).

NMR (IDG1, 1 4-0.5), -0.69, Le.23, 33.39, 34.11, 46.46, (7.84, 49.81, 115.96, 113.3), 126.90, 127.39, 123.46, 132.99, 137.30, 140.20, 140.81, 111.64, 142.05, 144.63.

Preparation of N--1,1-Dimethylethyl.-1,1-dimethyl-1- 1,3,6,7-tetrahydro--phenyl-s-indader-1-yl/dilanamine, dilithium sait.
N-(1,1-Dimethylethyl.-1,1-dimethyl-1--1,5,6,7-tetrahydro-3-phenyl-s-indaden-1-yl)silanamine (10.6551 g, 0.02047 moles) was othered in nexane (100 mb, as nBuli (0.070 moles, 35.00 mb of 2.0 M solution in dyclonexane) was added slowly. This mixture was then allowed to stir overnight during which time no salts drashed out of the dark red solution. After the reaction period the volatiles were removed and the residue quickly washed with nexane (0.00 mb). The dark red residue was then pumped dry and used without further purification or analysis (9.6517 q, 67.70 yield).

Preparation of Dichloro[N-(1,1-dimethylethyl)-1,1-dimethyl-1- $\{(1,2,3,4,5-\eta)-1,5,6,7-\text{tetranydro-}\}$ -phenyl-s-indacen-1-yl]silanaminato(2-)-N}titanium.

N-(1,1-Dimethyletnyl)-1,1-dimethyl-1-(1,5,6,7-tetrahydro-3-phenyl-s-indacen-1-yl)silanamine, dilithium salt (4.5355 g, 0.01214 moles) in THF (50 mL) was added dropwise to a slurry of TiCl₃(THF)₃ (4.5005 g, 0.01214 moles) in THF (100 mL). This mixture was allowed to stir for 2 hours. PbCl₂ (1.7136 g, 0.006162 moles) was then added and the mixture allowed to stir for an additional nour. After the reaction period the volatiles were removed and the residue extracted and filtered using toluene. Removal of the toluene resulted in the isolation of a dark residue. This residue was then slurried in nexane and cooled to) C. The desired product was then isolated via filtration as a red-prown crystalline solid 0.5280 g, 43.5% yield).

20

- B. DEBE - 1001 - 1 - 10.17 - 10.1

- ### MMR 2.0, a about 7, a mover 1, a mover 1, a mover 2, a move 2, a
- 77 NMR (JDL): (3.97, 3.87, 3.87, 3.80, 32.80, 32.74, 62.88, 78.14, 119.18, 121.97, 123.84, 127.18, 129.83, 129.37, 129.88, 138.34, 124.87, 138.34, 129.38, 129.37, 129.88, 138.37, 138.38, 138
- 15 Treparation of Dimethyl(N+01,1-dimethylethyl -1,1-dimethyl-1-(1,2,3,4,5-m -1,5,6,7-fetranydro-denyl-denyl-dendagen-1-yl)cilunaminato 2-denyl-ditanium.

Cichlorofth 1,1-simethylethyl =1,1-simethyleth 1,2,0,4,5-m = 1,5,6,7-tetranyaro-s-phenyl-l-indagen-l-yi)cilanaminato .- -50titanium

- 20 0.4970 ;, 0.301038 moles was ittired in diethylether 10 mL, as MeMdBr 0.0021 moles, 0.70 mL or 10. Misplution in diethylether; was added blowly. This mixture was then othered for 1 hour. After the reaction period the volatiles were removed and the residue extracted and filtered using hexane. Removal of the nexane resulted in the
- isolation of the desired product is a solden yellow solid $(0.4546\ \mathrm{g},\ 66.7\%\ \mathrm{yield})$.

H NMR (C_5D_6) : ± 0.071 s, 3 H), 1.49 (s, 3 H), 0.70 (s, 3 H), 0.73 (s, 3 H), 1.49 (s, 3 H), 1.7-1.8 (m, 2 H), 2.5-2.8 (m, 4 H), 6.41 (s, 1 H), 7.29 (t, C_5D_6): Hz, 2 H, 7.48 s, 1 H, 7.72 d, C_5D_6

- TO NMR CLOCK 12.13, 4.61, LT.12, T2.86, 33.00, .4.73, 58.66, 08.62, 118.62, 121.98, 124.26, 127.32, 128.63, 128.98, 131.23, 134.39, 136.38, 143.19, 144.85.
- 35 Cocatalyst (bis(hydrogenated-tallowalkyl)methylamine: (B-FABA) Preparation.

Methylcyclohexane (1200 mL) was placed in a 2L cylindrical flask. While stirring, bis(hydrodenated-tallowalkyl,methylamine (ARMEEN® M2HT, 104 d, ground to a granular form) was added to the

- flask and stirred until completely dissolved. Aqueous RCl 1M, 200 mL) was added to the flask, and the mixture was stirred for 30 minutes. A white precipitate formed immediately. At the end of this time, LiB(C,Ft), Et O 3 LiCl MW = -37.3; 177.4 % was added to the flask. The solution began to turn milky white. The flask was
- 45 equipped with a 6" Vigreux column topped with a distillation apparatus and the mixture was neated 140 °C external wall temperature). A

The Two-phase objection has now only accountly hazy. The mixture was allowed to find the term temperature, but the total the were placed in a 1 D separatory timber. The Equation was removed and the agreed, and the organizative was without twice with H and the aqueous layers again indicated. The normalized methylogical example to contain the were measured to contain the weighted nearly letter. Etc.:

The solution ACC mL) was transferred into a 1-1 flask, sparged incroughly with nitrogen, and transferred into the drypox. The solution was passed through a solumn 1" diameter, 6" height) dontaining 13% molecular sieves. This reduces the level of Et O from 0.48 wt percent to 0.23 wt percent. The material was then stirred over fresh 13% sieves 20 % for four hours. The Et C level was then measured to be 0.13 wt percent. The mixture was then stirred approximately 10 ppm. The mixture was filtered using a funnel equipped with a diass frit having a pore size of 10-15 µm to give a clear solution (the molecular sieves were rinsed with additional dry methylsycionexane). The concentration was measured by gravimetric analysis yielding a value of 16.7 wt percent.

Polymerization

Ethylene/styrene copolymers F-H were prepared in a 6 gallon (22.7 L), cil jacketed. Autoclave continuously stirred tank reactor (CSTR). A magnetically scupled aditator with Lightning A-320 impellers provided the mixing. The reactor ran liquid full at 475 psig (3,275 kPa). Process flow was in at the bottom and out of the top. A heat transfer cil was circulated through the jacket of the reactor to remove some of the heat of reaction. At the exit of the reactor was a micromotion flow meter that measured flow and solution density. All lines on the exit of the reactor were traced with 50 psi (344.7 kPa) steam and insulated.

Ethylbenzene solvent was supplied to the reactor at 30 psig (207 kPa). The feed to the reactor was measured by a Micro-Motion mass flow meter. A variable speed diaphragm pump controlled the feed rate. At the discharge of the solvent pump, a side stream was taken to provide fluon flows for the datalyst injection line. I lb-hr 1.45 kg/hr) and the reactor agitator 0.75 lb/hr 3.34 kg/hr). These flows were measured by differential pressure flow meters and controlled by manual adjustment of micro-flow needle valves.

Thinnibited Cyvrone manager was capped in the resonor at thinsis Difference of the constraints are measured by a Missi-Motion mass flow meter. A marriable object meaningm nump montrolled the toes rate. The onlyrene of seams was mined with the remaining colvent otream. Othywene was cumpiled to the reactor as 600 poly (1,137 Mpa). The Athylene tream was measured by a Midro-Motium mass of w meter just prior to the Research value controlling flow. A Brooke flow meter/controller was lied to deliver avarogen into the ethylene atream at the outlet of the ethniene control vilve. The ethniene hydrogen mixture combines with the solvent styrene orream an ambient temperature. The temperature of the solvent monomer as it enters the reactor was propped to 45 °C by an exchanger with -1°C plyoul on the jacket. This stream entered the sottom is the seastor. The three component satisfyot system and its solvent clash also entered the reactor at the notition but through a different fort than the monomer otream. Openaration to the datalyou components took place in an inert atmosphere flowe nox. The filuted simponents were put in hitrogen padded sylinders and sharged to the satalyst sun manks in the process area. From these run tanks the datalyst was pressured up with piston pumps and the ilow was measured with Micro-Motion mass flow meters. 20 These streams compane with each other and the datalyst flush solvent just prior to entry through a single injection line into the reactor. Polymerization was stopped with the addition of catalyst kill (water mixed with solvent) into the reactor product line after the micromotion flow meter measuring the solution density. Other polymer additives can be added with the datalyst kill. A static mixer in the line provided dispersion of the datalyst will and additives in the reactor effluent stream. This stream next entered post reactor neaters that provide additional energy for the solvent removal flash. This flash occurred as the effluent exited the post reactor heater and the pressure was dropped from 475 psig 3,275 kPa) down to ${\sim}250 \text{mm}$ of pressure absolute at the reactor pressure control valve. This flashed polymer entered a not bil jacketed devolatilizer. Approximately 85 percent of the volatiles were removed from the polymer in the devolatilizer. The volatiles exited the top of the devolatilizer. The stream was condensed with a glycol ploketed exchanger and entered the suction of a vacuum pump and was discharged to a glycol macket solvent and styrene/ethylene separation vessel. Colvent and styrene were removed from the bottom of the vessel and ethylene from the top. The ethylene stream was measured with a Midro-Motion mass flow meter and analyzed for composition. The measurement of vented ethylene plus a dalculation or the dissolved dasses in the solvent/styrene stream

were used to database one ethicients conversion. The paymer deparated in the devolutilizer was competed to with a dear number of LORE-30 sevolutilizant vacuum extruder. The dry polymer exit, the extruder as a single strand. This trand was cooled as it was pulled curbudh a categoristic, the extruder as categoristic, the extruder as categoristic, the extruder as categoristic, the extruder was cooled as it was pulled curbudh as categoristic pulled with it motive companies.

The actual preparation conditions for each cample are cummarized in Tuple 1.

Table 2

MMAO ³⁷ Fi Ratio	9.9
Co-B/T1	1.24
	B-FABA B-FABA
- Gatályst	ES 1
Conversion	E/5-G 86.2 28.27 2 48 17 10.5 92.3 ES 1 B-FABA 1.24 ES Catalyst is dimethyl [H-(1,1-dimethyl 11,1-dimethyl-1-[(1,2,3,4,5,1)]] 5 P. S. D. S
Hydrogen Styrene Flow Flow IF In Ib/hr	7 8 10 5 20 0 dimethyl-1-
Hydrogen Flow	5 5 ethy 1, 1
Sample Reactor Solvent Ethylene Prow Flow C lb/ht lb/ht E/S-F 91 11 40	2.48 1.0 1,1-dimethyl
Solvent Flow lb/ht	28.27 19.24 1methyl (N-
Reactor Solv Temp FIG	86.2 61.2 talyst is d
Sample	E/S-G E/S-H a ESI can

yllsilunaminatoi?-)-NJ- titanium. b CGC-7 catalyst is (t-butylamidoldimethyliteitemethylisy) begentadiane introdumenti) j. Jentadiane prepared as described in U S Patent 15,556,924, Example 17 c BFABA is bis-bydrogenated tallowally) methylummonium tetrakis (pentatluorophenyliteate) d a modified methylaluminoxune commercially varilable from Akzo Robel as HMAD-3A

ETHYLENE (TUBENE 1 7 1 L'OMER D. 1 . 1 . 1 . 1 personn) - 1. 1 male cordent sturene.

therpolymer had fine but wint interprets a weight person dryrene and more perfect that the but wint interprets a weight person dryrene along more perfect three belows density and the but mate 190°C, and Fig. along the set of insigning the Big. 1 in these transition remperature COO derive to a 100°C.

ETHYLENE/CTYRENS CYFOLYMER B (1.5 wt percent; 1...5 mole percent styrene)

36e General Procedure A. The resulting ethylene styrene interpolymer had the following properties: Weight percent styrene = \$1.8; mole percent styrene = \$2.5.

ETHYLENE/STYRENE COPOLYMER 1 04.1 At percent; 30.5 more percent styrene:

See General Procedure A. The resulting ethylene/styrene interpolymer had the following properties: weight percent styrene = 64.1; mole percent styrene = 32.5; sentity = .3982; melt flow rate (190°C, 2.2 kg) = 1.84; (lass transition temperature DSC derived) = -4.6°C.

20 ETHYLENE/STYRENE COPOLYMER D +67.0 wt percent; 35.4 mole percent styrene)

See General Procedure A. The resulting ethylene/styrene interpolymer had the collowing properties: weight percent styrene = 67.0; mole percent styrene = 65.4; density = 0.326; melt flow rate 190°C , 1.2 Kg) = 1.7; class transition temperature 000 terived) = -3°C .

ETHYLENE/STYRENE COPOLYMER E (72.5 wt percent; 41.6 mole percent styrene)

See General Procedure A. The resulting ethylene/styrene interpolymer had the following properties: weight percent styrene = 72.5; mole percent styrene = 41.6

ETHYLENE/STYRENE COPOLYMER F (28.8 wt percent; 4.82 more percent styrene)

Gee General Procedure B. The resulting ethylene Styrene interpolymer had the collowing properties: weight percent Styrene = 28.8; mole percent Styrene = 9.92; melt flow rate (190°C, 2.2 Kg) = 1.0. Further properties are shown in Table 2.

FINALUM (TUDEME , STICTURE) which represents Liber the terms of the terms of the second types.

lee pensone line insure of The issulting enoughere organie enterpring the control of the control

One seneral Problemure B. The resultand ethylene orgrene thaterpolymer had the fallowing properties: weight percent styrene = 73.3; move percent styrene = 42.1; mest sluw rate 190°n, 0.2 Eq; - 2.7. Surther properties are shown in Table 1.

TOLYCLEFTH A 1 - A NAME CONDITY (LIVETRY LONG AVAILABLE FROM THE LOW THEMSEL. - MEANY ALL OF 1 TON HAVING THE FULL WIND PROPERTIES: GENOLTY 1.4504 (Mr. melf fillw rate ().600 Toka (= 10.4)

POLYCLEFIN 3 to an orbytone detene appolymer available from The bow Inemical Company as COWLEX® CO474A having the following properties: density = 0.9173 from Finelt flow rate 0.200°C/5 kg/ = 0.20°C/5

BLOCK COPCLYMER A is VECTORIM (211-D available from Demos Polymers which is a Styrene-Isoprene-Styrene block topolymer containing 29.6 wt. percent styrene and 70.4 wt. percent isoprene having the following properties: melt flow rate (200°C, 5 kg) of 10.6 g.10 min.

BLOCK COPOLYMER B is VECTORIM 9008-10 available from Dexico Polymers which is a Joyrene-Butadiene-Styrene block sopolymer sontaining 18.5 wt. percent styrene and 71.8 wt. percent isoprene having the following properties: melt flow rate .200°C 5 kg; or 12.2 g/10 min.

BLOCK COPOLYMER C is VECTORIM T400-D available from Dexco Polymers which is a Styrene-Butadiene-Styrene block copolymer with a molar ratio styrene-butadiene of 31.8768.2 having the following properties:

melt flow rate .200°C 5 kg; wi 17.3 g/10 min. and an oil content of 33.3 percent by weight.

BLOCK COPOLYMER D is KPATONIM G 1652 available from Shell Chemical Company which is a styrene-ethylene/butylene-styrene triblock copolymer containing of who percent styrene and TI who percent athylene/butylene.

EMAMPLES 1 - - AND 1 MEARATINE SMEER IMENTS A -

An expension of the second of the control of the appropriate to

A compression monotoner present (181710 sector823) 1-X-MS-X24 was theheares is continuous \mathbb{R}^{n+1} . We have a few consisting is stainless thed parking that is installed TDE crafted alaborism oneets, and a metal mase i approximately of mm him by 171 mm wide by 1.13 mm thick was prepared. Into the chase was evenly opread an amount or Block Supelymer neclets Simponent A indicated in Table . From a slab of Jumponent B was out an amount of polymer indicated in Table I which was evenly instributed in the chase on top of the block polymer. A layered structure with TFE doated sheets was formed next to the polymer. The sundwich was preheated in the press at hear minimum preseure for approximately one minute, then pressure of the press was increased to about U., 60 Mu of ram torce for 10 seconds. The imase and the cused blend was removed when dooled to sufficient rigidity. The still warm blend was placed into an operating roll mil-(155°C) for a minutes total. The blend was temporarily removed, folded, and replaced on the muli at least twice. The plend was removed from the roll mill and allowed to cool. This material was used for further fabrication. 20

B. Preparation of Test Jample.

A compression molding press—PHI Model CB234 T-M-MS-X24) was preheated to CCO°C. A 7.) gram portion of the blend from A above was evenly distributed into a 75 mm wide by 115 mm high by 0.34 mm thick chase layered between TFE-coated glass sheets and metal backing plates. The following temperature program was followed: preheat 3.0 minutes at minimal pressure; medium force 0.5 minutes /10,000 kg ram force); material cure 3.0 minutes /20,000 kg ram force).

C. Preparation of Test Cample.

A compression molded circle was prepared from another portion of the roll-milled blanket prepared in A above in a manner similar to the first sneet prepared in B above except that 14.0 grams of blend are used in a chase which has "6 mm diameter similes and which was approximately i.1 mm thick. Tircles are out in half and soupled to required intexhess for those A testing.

The Therman of the property of the service

The filat specimens are overex in 1807 for it nours or more prior propagator in 8 above juing a dinterial Triorimanus solorimeter CCSP-4

stimilar to ASTM D (10.8). Tensilo, modulus, and ast operamens are die crotopsi was 10. persont silvantion, and objective det ceconds, measure deroent sensile set, settined as dinal dauge-initial dauge on 100 and and a sensite set and a sensite det cause set and conditions dauge-initial dauge of the cetween graphs of the cetween graphs as when corps persones apparent sites removing slack.

Shore A hardness was determined by ASTM D 0240.

Force at 300 percent elongation (300 percent modulus) was determined by ASTM $2.112\,\mathrm{L}_\odot$

Fercent relaxation was a defined in ADTM 0 2/91-94 after extension to 10% percent lundation and to based in corpe value upon reaching maximum elongation and apter 10 seconds.

The test results are provided in Table 3.

Table 3

			Table	⊋ 3				
DI	Ex. 1	Ex. 2	Ex. 3	Ex. 4	Ex. 5	⊏X. છે	Ex. 7	=x. 8
Blena#	6252-5-8	6252-5-3		6252-5-20	6252-5-4	6252-5-9	 6252-5-15	6252-5-13
Component B	E/S Dooryme	E/S 2000vmer	⇒polymey 	. ∑ES	ES	£3	E/S	ES
Туре	. A	A	A	A	Cipalymer B	_ Cipovimer B	. ⊝polymer B	- Cooolymer
wt. % ethylene	55	55	55	55	48	48	48	48
wt. % styrene	45	45	45	45	52	52	52	52
Amount, grams	8	9	19	19	9	8	8	8
wt. %	20	20	50	50	20	20	20	$-\frac{3}{20}$
Component A	Blook Copatymer	Block Copolymer	Block Coporymer	Block Copatymer	Broak Coccivmer	Blook	3/ock	Biook
Туре			В	C	A	Coccinmer -	Copolymer	Copolymer
Amount, grams	32	36	19	19	36	32	32	D 24
wt. %	80	80	50	50	80	80	$-\frac{32}{80}$	31 80
% Haze	23	17	53	60	28	33	65	98
Shore A Hardness	65	59	62	53	54	63	47	73
Ultimate Tensile								
psi	4,398	3,163	3,147	2.560	3,437	3,978	1,603	5 154
kPa	30,323	21,808		17.651	23.697	27.427	11.052	
% Elongation at	- 				20.007	21.721	11,002	35,536
Break	850	925	542	758	1,017	858	1,092	558
Force at 300% Bongation(300% Modulus)								
psi	635	507	640	205	12.1			
kPa	4,378	597 4,116	648 4.468	395 2. 723	434 2.992	445 3,068	597	846
	,			2.720		3,000	4,116	5.833
% Relaxation (after 150% elongation)	22	21	21	19	20	22	21	17
% Set (after 150% Hongation)	· 11	4.4	12	!		_	 :	
- 34401)		11	13	13	10	12	13	11

Table 3 contd.

		Lau	ole 3 :	contd.				
	B C 9	Ex. 10	± 11	⇒c 12	Ex. 13	Ex. 14	Ex 15	Ex. 16
∃ena #	6252-5-5	6252-5-10	6252-5-	32 52- 5-	 625 2-5- 6	6252-5-		6252-5
	ES		21	22		11	16	14
Component B	Capolymer	_Spotymer	ES S	ES COCOMMO	ES.	<u> </u>	ē\$	₽S
Type	C	C	C	C	Capaymer	Copowmer	Coporymer	
wt. %ethylene	36	36	36 ~	· 36 · ·	3 3	33	33	
wt. %styrene	64	64	64	. 64	. 67	67	67	33
Amount, grams	8	8	19	19	· 8 –	8	8	67
wt. %	20	20	50	50	20	20	<u></u>	8
Component A	Bock	Bock	Воск	Block	- 			20
	Capalymer			Copolymer		Bock	Book	Воск
lype	A	В	В	C	- - -	В	C	D
Amount, grams	32	32	19	19	32	32	31	
wt. %	- 80	80	50	50	80	30		32
								80
%Haze	73	89		99	78	84	-00	
							99	99
Shore A Hardness	59	63	47	46	60	CC		
Utimate iensie				40	- 00	66	45	77
pa	2,958	3,745	819	729	2.875	2 700	1 707	
kPa	20.395	25,821	5,647	5.026	19.822	3,722	1,727	5,627
%Bongation at			3,047	J.020	19.022	20,002	11,907	38,797
Break	900	742	1.000	1.050	OFR	700		
Force at 300%		172	1,000	1,030	958	733	983 .	468
Bongation(300%								
Modulus)								
pg	466	648	247	255	040			
kPa	3,213	4,468	1,703	255	613	716	293	1,197
	J.2 15		1,703	1.758	4.226	4.937	2.020	8.253
%Relaxation (after							1	
50%elongation)	20	22	22	20			ł	
			33	33	24	23	18	18
%Set (after 150%								
elongation)	12	10	40	4=			!	
- 3-0011)	14	10	18	17	12	12	17	10

Table 3 contd.

			.e 3 cont	 .			
	Ex 17	Ex 18	Ex. 19	Comp. Expt. A*	Comp. Exat. B*	Comp. Expt. C*	Comp. Expt. D*
Bend#	6252-5 -5	· 252-5-1 0	52 52 -5-21	32 52 -5-22	92 52- 56	6252-5-11	62 52-5 -16
Comparent B	5SCapatymer	t:SCaparvmer	#S Capatymer			avaletin	Patyalefin
Туре	Ε	E	Ε	Α	А	B	В
wt. % ethylene	27	27	27		· · · · · · · · · · · · · · · · · · ·		
w.t. %styrene	73	73	73	<u></u>			
Amount, grams	8	8	19	19	20	19	20
wt. %	20	20	 50	50	50 -	50	50
Component A	Block Capatymer	Block Cuppatymen	Block Capatymer	Block Capolymer	Block Capatymer	Block Capatymer	Block
Туре	Α	B	В	В	C	В	С
Amount, grams	32	32	19	19	20	19	20
wt %	. 80	80	50	50	50	50	50
%Haze	<u>-</u>	83	99	94	93	99	<u> </u>
Share A Hardness	61	66	87	87	- 33 - 89	35 86	99
Utimate Tensile			•	0/	ω-	ω	85
bai	3,628	3,981	2,679	3,171	2,797	2,791	2545
kРа	25,014	27,448	18,471	21,863	19,285	19,243	2,545
% Bongation at Break	958	783	417	842	750	742	17,547 717
Force at 300% Bongation(300% Modulus)					100	7-12	717
ba	737	814	1,856	1,736	1.693	1,192	1,180
kPa	5.081	5,612	12,797	11,969	11,673	8,219	8,136
% Relaxation (after 150 alongation)	18	21	59	45	45	30	29
% Set (after 150% elongation)	11	12	62	ස		40	39

^{*} Not an example or the present invention.

EXAMPLES 20 + 26

5 a) Compression molding:

Samples were melted at 190 °C for 3 min and compression molded at 190°C under 10,00° lb or pressure for another ... min. Subsequently, the molten materials were quenched in a press equilibrated at room temperature.

10 <u>ti Differential Canning Calbrimetry (DSC):</u>

A Du Pont 000-1010 was used to measure the thermal transition temperatures and heat if transition for the samples. In order to

eliminate previous thormal history, damples were first heated to about 160 °C. deating and obtains surves were descrated at 10 °C min. Welting t_m from descend neath and prystallization t₀ temperatures were resorded from the seak temperatures it the endotherm and exotherm, respectively.

Cynamic Mechanical Spectroscopy CMS::

Dynamic mechanical data were penerated using a Pheometrics RSA-II solid state analyzer, and melt pressed of 10 mil thick/ film test specimens. The DMS measurements were conducted at a step rate of 5 %0 min and a fixed frequency of 10 rad/sec. The class transition temperature Tg) of the samples was determined from the tan ô peak maximum.

3: Shear Pheology:

Discillatory shear rheology measurements were performed with a Sheometrics RMS-400 rheometer. Melt rheological properties were monitored at an isothermal set temperature if 199°C in a trequency sweep mode, using parallel plate test geometry.

e) Mechanical Testing:

Tensile properties of the compression molded samples were measured using an Instron 1145 tensile machine equipped with an extensiometer. ASTM-D638 samples were tested at a strain rate of 5 min⁻¹. Micro-tensile samples were tested at a speed of 5 in/min at - 10° C. The Youngs Modulus (E, Mpa) measured was the average of four tensile measurements was given. The standard deviation for the altimate properties was typically about 10° 3 of the reported average value. Also measured was the yield stress at the inflection point of the stress strain curve σ_{φ} , Mpa) and the Ultimate Tensile stress at preak $(\sigma_{b}$, Mpa).

f) Tensile stress relaxation:

Uniaxial tensile stress relaxation was evaluated using an Instron 1145 tensile machine. Compression molded film (~ 20 mil thick) with a 10 mil gauge length was deformed to a strain level of 50% at a strain rate of 20 min $^{-1}$. The force required to maintain 50% elongation was monitored for 10 min. The magnitude of the stress relaxation was defined as Sr, the percentage = $+f_1 - f_f / f_1$) x 100 where f_1 was the initial force and f_f was the final force.

g) Thermal Mechanical Analysis.

Upper service temperature (TMA(lmm)) was determined from a thermal mechanical analyzer (Perkin Elmer TMA 7 series) scanned at 5

Dimin and a load of 1 News no and betined as the point at which the prope penetrates 1 mm into the sample.

Example 10.

Example 10 was a plend containing 75 % by weight of ESI # F and 25 % by weight of Block Cupolymer J. The blend was prepared by blending in a Hauke Freemix 3000 bowl mixer. The dapacity of this mixer was 310 cc. ptimum volume for offective mixing was approximately 76% or 220 st. Salculations were made considering mensity and constituency or each component to prepare a dry blend of the materials to achieve a 70% volume fill. The dry blended materials were then added stepwise into the preheated calibrated bowl as the rotors were turning at 30 rpm. The materials were heated to approximately 250 degrees Centigrade. After a small melt heel was established in the mixer, small increments of dry blend were added and allowed to meit and incorporate into the heel before more blend was added. This continued for approximately two minutes until all the blend was added. A sealing ram was then lowered on to the melt bowl and the melted blend was allowed to mix by roller brade action for an additional ten minutes. At the end of this time the rotors were stopped, the mixer was dismantled, and the melt blend was removed and allowed to cool for further testing and analysis.

Example 21.

Example 21 was a plend containing 25 % by weight of ESI # F and 75 % by weight of Block Copolymer D. The blend was prepared essentially as for Example 20.

Example 22.

Example 22 was a plend containing 75 % by weight of ESI # G and 25 % by weight of Block Copolymer D. The blend was prepared essentially as for Example 20.

Example Lo.

Example 13 was a plend containing 5. by weight to ESI # 3 and 50 by weight of Block Copolymer C. The plend was prepared assentially as for Example 20.

Example 14.

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Example 24 was a plend containing 18 % by weight of ESI # G and 75 % by weight of 31cox Copolymer 3. The plend was prepared -ssentially as for Example 23.

Example 15.

Example 15 was a plend containing 75% by weight of ESI # H and 25% by weight of Block Copolymer D. The blend was prepared essentially as for Example 20.

Example 26.

Example 25 was a plend containing 25 % by weight of ESI # H and 75 % by weight of Block Copolymer D. The plend was prepared essentially as for Example 20.

These data lilustrate that as the mole of styrene content of the ESI component of the blend increases then the sistress relaxation also increases. These data also illustrate that below approximately 22 mole styrene content of the ESI component of the blend, as the styrene content of the ESI component of the blend, as the styrene content of the ESI component of the blend decreases the Ultimate tensile value increases, whereas above approximately 22 mole styrene content of the ESI component of the blend, as the styrene content of the ESI component of the blend, as the styrene content of the ESI component of the blend, as the styrene content of the ESI component of the blend increases, the Ultimate tensile value increases

Table 4

Blend (Blend Components	12	110/12	w L *	mol &S.	wt t mol & S* Tq (DSC)	7.11	Taibisi	T. 1 1 14	1111121
				à P.S		ບ			0 0 0	
E/: F	E:329	1.0	7.7	3.3	5.82	-17.6	73.3	(i .	!
E/S-C	ES52	1:1	7.6	1.0	+	-11.9	,	-1-		
E/S·H	ES75	2.2	9.2	8. E	42.50	20.6		30.6		
Block.							10.3	-13.4	101	
cobol. D										
Blends	(onponents		Compostn					-		
:			(Wt 8)							
Ex 20	E/S-F/BC D		75/25			-17.6	77.6	-46.7	-5.3	
Ex 21	E/S-F/BC-D		25/75				18.9/7	-44.5	-7.0	9.66
							9.3			,
Ex 22	E/S-G/BC-D		75/25			-11.6		-45.1	3.0	
Ex 23	E/S-G/BC-D		50/50			-10.3	15.8	-44.5	-3.0	1001
Ex 24	E/S-G/BC-D		25/75					-44.5	2 3	101
Ex 25	E/S-H/BC-D		75/25			20.1		-42.3	30.0	7.101
Ex 26	E/S-H/BC-D		25/75					-43.9	31.2	100 7
									1.10	

(Wt% styrene/104)/((Wt% styrene/104) + (Wt% ethylenergas)

Table 4 contd.

	Stress	Relax, e		3.	-	00.7	12.4	20.3		:			30.8		2	77	.1.4	19. 2	, ,	0.75
	Ult Ten, Si			3.54	-			9.05	_				32.2	1	F . F.7	-	5.3 	19.4	+	-
			_ <u> </u>	_		1										-			 	-
	Eich)		1	7	058		2	(4)		-			C & 5	300		0	17.	2.5	113	7
	Flex	Mod, MFa		02.4			ļ	30.5					43.9	33 4	- C		F . 4	41.9	104.1	2 CZ
	Shore A		00/00	50/06	62/29	10/96	10.00	81/80				30,00	86/85	83/82	68/64	14/77	7//5/	76/74	96/93	82/81
	TMA C		a a		20	99	23	109				000	0	97	63	98	0	ص ص	71	93
	& Xtyl		3									1,60	0.01							
										Compostn	(×t %)	75/25	77.67	25/75	75/25	50/50	36/36	67/67	75/25	25/75
	alend components		ES29	6303	7007	ES75			24.000000	components		E/S-F/BC-D		E/2-1/BC-D	E/S-G/BC-D	E/S-G/BC-D	F/8-6/BC-D	2 2 2 2 2 2	E/S-H/BC-D	E/S-H/BC-D
0 7-010	on puerd		E/S-E	F/8-C	0.073	E/S-H	Bleck	Copol. D	-			Ex 20	1	EX 21	Ex 22	Ex 23	Ex 24	1	<u>_</u>	Ex 26

WHAT IS CLAIMED IS

... A thermoplastic elastemente clend composition :cmpricing: The trom of the lower mit by roem of a cityrenic block sopolymer; and 5 from 1 to 30 weight percent it an interpolymer B of (1) at least one aliphatic lpha-blefin and (1) at least one vinylidene •3 aromatic monomer or 3 a compination of at least one vinylidene aromatic monomer and at least one hindered aliphatic vinylidene 8 monomer, wherein said vinylidene aromatic monomer is present in an q amount of from 0.5 to 15 mole percent or from 17 to ${\mathfrak n}{\mathfrak h}$ mole percent in . . said interpolymer.

12

- 1 1. The composition of Claim 1 wherein the elastomer block copolymer component A) comprises at least one segment of a styrene polymer and at least one segment of the following: isoprene polymer, 3 butadiene polymer, isoprene/butadiene copolymer, ethylene/butylene copolymer or an ethylene/propylene copolymer; and component (B) is an 5 6 interpolymer of ethylene/styrene.
- 1 3. The composition of Claim 1 wherein the styrenic block copolymer component (A) comprises a polystyrene-polyisoprenepolystyrene or polystyrene-polybutadiene-polystyrene; and component 3 B) is an interpolymer if ethylene/styrene.

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- The composition of Claim 1 wherein the styrenic block copolymer component A) contains from 10 to 35 weight percent of styrenic polymer segments and from 90 to 65 weight percent of elastomeric saturated clefin polymer segments, based on the total weight of the block copolymer.
- 1 5. The composition of Claim 1 wherein the styrenic block copolymer component (A) contains from 10 to 35 weight percent of 2 styrenic polymer segments and from 90 to 65 weight percent of elastomeric insaturated polydiene segments, cased on the total weight of the block appolymer.

P. The composition of Tlaim . wherein component B has a and heat of tusion of 1938 than of Jog. . The composition or claim 3 having a percent tensile set after .", reroent -lungation hr II percent in less and a 3 transmission percent hade to All percent or less and haze value being istained in 1.24 mm thirm sample. o. A fabricated article made from the composition of Claim 1. 3. A fabricated article made from the composition of 2 Claim 2. 13. A fabricated article made from the composition of Claim 3. 11. A fabricated article made from the composition of 2 Claim 4. 12. A fabricated article made from the composition of 2 Claim 5. 1 13. A fabricated article made from the composition of Claim 6. 14. A fabricated article made from the composition of Claim ~.

INTERNATIONAL SEARCH REPORT

PCT/US 37/18773

A CLA	SSIFICATION OF SUBJECT MATTER		
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(75) Inventors/Applicants (for US only): ESNEAULT, Calvin, P. 1 Published [US/US]: 14776 Colonel Allen Court, Baton Rouge, LA 70816 (US). EDMONSON, Morris, S. [US/US], Route 130 Mohawk Drive, Alvin, TX 77511 (US), CHEUNG, a Yunwa, W. [US/US]: Apartment 810, 510 That Way, Lake 1 Jackson, TX 77566 (US), GUEST, Martin, J. [GB/US], 106 Greenbriai, Lake Jackson, TX 77566 (US), HO, Thoi, H. [US/US]; 54 Orchid Court, Lake Jackson, TX 77566 (US).

With international search report Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of imenaments.

(54) Title: BLENDS OF ELASTOMER BLOCK COPOLYMER AND ALIPHATIC ALPHA-OLEFIN/MONOVINYLIDENE ARO-MATIC MONOMER AND/OR HINDERED ALIPHATIC VINYLIDENE MONOMER INTERPOLYMER

(57) Abstract

A thermoplastic elastomene blend composition comprising: (A) from 99 to 1 weight percent of a styrenic block copolymer; and (B) from 1 to 99 weight percent of an interpolymer of (1) at least one aliphatic α -olefin and (2) at least one vinylidene aromatic monomer or (3) a combination of at least one vinylidene aromatic monomer and at least one hindered alibnatic vinylidene monomer, wherein said vinylidene aromatic monomer is present in an amount of from 0.5 to 15 mole % or from 17 to 65 mole % in said interpolymer, and fabricated articles made from such composition.

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